Mapping Region-Specific Longitudinal Cortical Surface Expansion from Birth to 2 Years of Age

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The human cerebral cortex develops rapidly and dynamically in the first 2 years of life. It has been shown that cortical surface expansion from term infant to adult is highly nonuniform in a cross-sectional study. However, little is known about the longitudinal cortical surface expansion during early postnatal stages. In this article, we generate the first longitudinal surface-based atlases of human cortical structures at 0, 1, and 2 years of age from 73 healthy subjects. On the basis of the surface-based atlases, we study the longitudinal cortical surface expansion in the first 2 years of life and find that cortical surface expansion is age related and region specific. In the first year, cortical surface expands dramatically, with an average expansion of 1.80 times. In particular, regions of superior and medial temporal, superior parietal, medial orbitofrontal, lateral anterior prefrontal, occipital cortices, and postcentral gyrus expand relatively larger than other regions. In the second year, cortical surface still expands substantially, with an average expansion of 1.20 times. In particular, regions of superior and middle frontal, orbitofrontal, inferior temporal, inferior parietal, and superior parietal cortices expand relatively larger than other regions. These region-specific patterns of cortical surface expansion are related to cognitive and functional development at these stages.

Keywords: cortical surface expansion, longitudinal cortical development, longitudinal surface-based atlas, infant cortical folding

Introduction

The adult human cerebral cortex is a highly convoluted and variable structure composed of sulci and gyri (Ono et al. 1990), which emerge in late gestation (Retzius 1890; Chi et al. 1977; Dubois et al. 2008). At term birth, all major sulci and gyri have been well established (Chi et al. 1977), although both brain volume and cortical surface area are only one-third of those of adults (Thompson et al. 2007; Hill, Dierker et al. 2010). After birth, tertiary folding structures are still undergoing development (Chi et al. 1977; Ono et al. 1990; Armstrong et al. 1995; Toro and Burnod 2005; Nie, Li et al. 2011), in concert with the rapid development of related cognitive and motor functions (Gilmore et al. 2007). Understanding normal cortical surface development in the early postnatal period would provide important insight into neurodevelopmental disorders (Gilmore et al. 2007).

Owing to the cellular nonuniformities, it is anticipated that there would be regional differences in macroscopic aspects of postnatal cortical development (Hill, Inder et al. 2010). Hill et al. (Hill, Inder et al. 2010) generated the first surface-based atlas of cortical structures of term-born infants using a small population of 12 healthy subjects. On the basis of the atlas, they compared the cortical surfaces of the 12 infants and 12 adults cross-sectionally to demonstrate the highly nonuniform postnatal cortical surface expansion, with the lateral temporal, parietal, and frontal regions expanding nearly twice as much as other regions in the insular and medial occipital cortex (Hill, Inder et al. 2010). However, cross-sectional studies are likely affected by variations of intersubjects and cohort effects (Gogtay et al. 2004), and to date the longitudinal cortical surface expansion in the early postnatal stage still remains largely unknown. In this article, we constructed the first longitudinal surface-based atlases of human cortical structures at birth, 1 year of age, and 2 year of age from a population of 73 healthy subjects. The surface-based atlases were generated by the landmark-free groupwise registration of cortical surfaces; thus, the generated atlas is unbiased to any individual and can also capture the variability of cortical folding. We then used the longitudinal surface-based atlases to study the longitudinal cortical surface expansion in the first 2 years of life. We find that cortical surface expansion is both age related and region specific. In the first year, the cortical surface area expands dramatically, with an average expansion of 1.80 times. Specifically, regions of superior and medial temporal, superior parietal, medial orbitofrontal, lateral anterior prefrontal cortices, postcentral gyri, and occipital lobes on both hemispheres expand relatively larger than other cortical regions. In the second year, cortical surface area expansion is still substantial, with an average expansion of 1.20 times. Regions of superior and middle frontal, orbitofrontal, inferior temporal, inferior parietal, and superior parietal cortices in both hemispheres expand relatively larger than other cortical regions. The region-specific patterns of cortical surface area expansion in the first and the second year are related to the cognitive and functional development at different stages.

Materials and Methods

Subjects

This study was approved by the Institutional Review Board of the University of North Carolina (UNC) School of Medicine. The parents were recruited during the second trimester of pregnancy from the UNC hospitals and written informed consent forms were obtained from all the parents. The presence of abnormalities on fetal ultrasound, or major medical or psychotic illness in the mother, was taken as exclusion criteria. The infants were free of congenital anomalies, metabolic disease, and focal lesions. None of the subjects was sedated for MRI. Before the subjects were imaged, they were fed, swaddled, and fitted with ear protection (Gilmore et al. 2011).

Complete 0-1-2 data of 73 normal infants was acquired in 30 singleton (20 males/10 females) and 43 twins (22 males/21 females; 7 monozygotic twin pairs, 10 dizygotic pairs, and 8 “single” twins). The mean gestational age at birth was 37.9 ± 1.6 weeks. The mean ages at the scan are 25.5 ± 10.8 days, 392.8 ± 22.1 days, and 758 ± 38.1 days, respectively. This same dataset has been used in a prior study of development of cortical gray matter volume (Gilmore et al. 2011).
MR Image Acquisition

Images were acquired on a Siemens head-only 3T scanner (Allegra, Siemens Medical System, Erlangen, Germany) with a circular polarized head coil. For T1-weighted images, 160 sagittal slices were obtained by using the 3D magnetization-prepared rapid gradient echo sequence: TR = 1900 ms, TE = 4.38 ms, inversion time = 1100 ms, Flip Angle = 7°, and resolution = 1 × 1 × 1 mm³. For T2-weighted images, 70 transverse slices were acquired with turbo spin-echo sequences: TR = 7380 ms, TE = 119 ms, Flip Angle = 150°, and resolution = 1.25 × 1.25 × 1.95 mm³ (Gilmore et al. 2011). Data were collected longitudinally at 3 age groups: neonates, 1 year of age, and 2 year of age. The data with motion artifacts was discarded and a rescan was made when possible. The variation of age at MRI for each scan is relatively small, and the population can be divided into age groups concentrated around 0, 1, and 2 years of age.

Image Processing

Before tissue segmentation and cortical surface reconstruction, all images were preprocessed using the following procedure. Noncerebral tissues were stripped by (Shi et al. 2012), followed by manual editing with ITK-SNAP software (Yushkevich et al. 2006) to ensure accurate skull removal. Cerebellum and brain stem are removed by in-house developed tools. Intensity inhomogeneity correction was performed with N3 method (Sled et al. 1998) to alleviate the influence on subsequent tissue segmentation. All images at each age were rigidly aligned to the age-matched infant image intensity atlas (Shi et al. 2011). Because of the low tissue contrast and difference in cortical shapes compared with adults, tissue segmentation methods developed for adult brain MR images do not work well on the infant brain MR images (Hill, Dierker et al. 2010). Therefore, tissue segmentation of infant brain MR images were performed by a dedicated longitudinally guided level-set–based method (Wang et al. 2011) by combining local intensity information, atlas spatial prior, cortical thickness constraint, and longitudinal information by longitudinal image registration (Shen and Davatzikos 2004; Xue et al. 2006) into a variational framework. Figure 1.1a,b,c shows a T2-weighted MR image at birth, corresponding skull-stripping result, and tissue segmentation result, respectively. After tissue segmentation, the lateral ventricles and subcortical structures were masked and filled, and then each brain was separated into left and right hemispheres as shown in Figure 1.1(d).

Cortical Surface Reconstruction

The inner cortical surface (the interface between white matter and gray matter) was obtained by correcting the topological defects (Fischl et al. 2001; Shattuck and Leahy 2002) and tessellating the white matter as a triangular mesh to ensure a spherical topology for each hemisphere. Since the transient subplate zone, which is interposed between the immature cortical plate and white matter, might still exist after birth, the inner cortical surface at birth was defined as the interface between the cortex plate and white matter zone (including white matter and transient subplate zone) (Nie, Li et al. 2011). The inner cortical surface was then deformed by preserving its initial topology for reconstruction of the central cortical surface (Li et al. 2012), which was defined as the layer lying in the geometric center of the cortex and approximately corresponding to the cytoarchitectonic layer 4 (Xu et al. 1999; Van Essen 2005; Liu et al. 2008). The central cortical surface was adopted for performing analysis of the cortical surface expansion as did in (Hill, Inder et al. 2010), since it provided a more balanced representation of gyral and sulcal regions (Van Essen 2005). The inner cortical surface, which had vertex-to-vertex correspondences with the central cortical surface, was further smoothed, inflated, and mapped to a standard sphere by minimizing the metric distortion between the cortical and the spherical representation (Fischl, Sereno, Tootell et al. 1999). Figure 1.1e,f,g,h shows an example of the reconstructed inner cortical surface, central cortical surface, the inflated inner cortical surface, and the spherical representation of the inner cortical surface, respectively. Note that the inflated cortical surface and the spherical representation are color-coded by the average convexity of the cortical surface, which integrates the normal movement of a vertex during cortical surface inflation and reflects the large-scale geometry of the cortical surface (Fischl, Sereno, Tootell et al. 1999). Figures 1.2 shows the longitudinal inner cortical surfaces (color-coded by mean curvatures) of a typical subject at 0, 1, and 2 years of age. Generally, gyri have negative mean curvatures, and sulci have positive mean curvatures (Note that we here adopt the inward-oriented normal vector field). As we can see, the major cortical folding of sulci and gyri has been well developed at term birth, and is well preserved during the cortical development from 0 to 2 years of age.

Surface-Based Atlas Generation

To build the longitudinal surface-based atlases of cortical structures, we need to determine the longitudinal vertex-to-vertex correspondences of cortical surfaces at different ages of each subject and also the cross-sectional vertex-to-vertex correspondences of cortical surfaces between different subjects at each age. To establish longitudinal vertex-to-vertex correspondences of cortical surfaces at different ages, for each hemisphere of each subject, the cortical surface at 0-year of age was registered to the corresponding cortical surface at 1-year of age using Spherical Demons registration method (Yeo et al. 2010). Spherical Demons aligns the cortical folding patterns mapped in the spherical space based on several geometric features of cortical surfaces, including mean curvature of the inflated cortical surface, average convexity of the cortical surface, and mean curvature of the cortical surface. It has been shown that the Spherical Demons achieved similar registration accuracy as FreeSurfer, but with much faster computational speed (Yeo et al. 2010). Similarly, the cortical surface of each hemisphere of each subject at 1-year of age was registered to the corresponding cortical surface at 2 years of age. The deformation map from 0 to 2 years of age of each hemisphere of each subject was obtained by concatenating the 2 deformation maps from 0- to 1-year of age and from 1 to 2 years of age by using the method in (Van Essen et al. 2011). In the next step, we focus on estimating atlases at different ages, and then build their relative deformation maps. The surface-based atlas of cortical structures at each age of each hemisphere was constructed by groupwise registration of cortical surfaces of all subjects at the age using Spherical Demons. Thus, the surface-based atlases are unbiased to any individual and can capture the variability of cortical folding. Specifically, at each age, first, an atlas consisting of the mean and variance of cortical geometry of all subjects was computed; second, all cortical surfaces were registered to the computed atlas; and finally, the atlas was updated based on the current registration results. The registration and atlas updating were iteratively performed. In our application, 3 iterations of registration were performed and reasonable results were achieved. On the other hand, to compute the deformation map between any 2 atlases (at 2 different ages), the longitudinal deformation of each subject project to the corresponding vertices of the atlas at the i-th year(s) (i, j ∈ 0, 1, 2, i ≠ j) was first obtained by concatenating 3 deformation maps: deformation from the atlas at the i-th year to the cortical surface at the j-th year, longitudinal deformation from the cortical surface at the j-th year to the cortical surface at the j-th year(s), and deformation from the cortical surface at the j-th year(s) to the atlas at the j-th year (s). The deformation map between any 2 atlases was thus obtained by averaging longitudinal deformations projected in the 2 atlas spaces of all subjects. Figure 1.3 illustrates the flowchart of generating longitudinal surface-based atlases of cortical structures at 0, 1, and 2 years of age.

Computing Cortical Surface Area Expansion

After building the longitudinal surface-based atlases, the central cortical surface of each hemisphere of each subject at 2 years of age was resampled to a standard-mesh tessellation with 163 842 vertices based on deformation from the surface-based atlas at 2 years of age to the individual cortical surface, thus establishing the vertex-to-vertex correspondences at each hemisphere across all subjects at 2 years of age. Based on the longitudinal registration result of each subject, the standard-mesh tessellation of each subject at 2 years of age was warped to 1-year of age and 0-year of age, respectively, thereby establishing the longitudinal vertex-to-vertex correspondences of each
subject. Note that the occupied area of each vertex was computed as one-third the sum of areas of all triangles associated with that vertex. And the fractional area of each vertex was computed as the occupied area of each vertex as the fraction of the total surface area. The cortical surface area expansion map from the \(i\)th to the \(j\)th year(s) of each hemisphere of each subject was computed by comparing the occupied vertex area at each vertex at the \(j\)th year(s) to that of its corresponding vertex at the \(i\)th year. The surface areal expansion map of each hemisphere of each subject was smoothed by using an iterative nearest neighbor-averaging procedure.

*Figure 1.* Illustration of cortical surface reconstruction and longitudinal surface-based atlases generation.

\(i, j \in \{0, 1, 2, i < j\}\) of each hemisphere of each subject was computed by comparing the occupied vertex area at each vertex at the \(j\)th year(s) to that of its corresponding vertex at the \(i\)th year. The surface areal expansion map of each hemisphere of each subject was smoothed by using an iterative nearest neighbor-averaging procedure.
on the surface mesh (Han et al. 2006). All smoothed surface area expansion maps were generated by 40 iterations of smoothing, approximately corresponding to 5 mm Gaussian smoothing kernel (Han et al. 2006). The smoothed surface area expansion maps of all subjects are averaged to generate the final surface area expansion map.

Results

Global Cortical Surface Area Expansion in the First 2 Years

Figure 2 shows the distribution of cortical surface areas of the 73 subjects at 0, 1, and 2 years of age, and Table 1 provides the mean and standard deviation of cortical surface areas of the 73 subjects. Note that all the cortical surface areas were measured in their native space. The cortical surface area of male is generally larger than that of female at each age, consistent to the findings with the brain volume of male being larger than that of female (Shi et al. 2011). Specifically, the cortical surface area is about 14%, 11%, and 12% larger in males compared with females at 0, 1, and 2 years of age, respectively, consistent to the findings that the cortical surface area is about 10% larger in males compared with females in adults (Van Essen et al. 2011). However, this is different from another study with a smaller sample size, which found no gender difference in infants (Hill, Dierker et al. 2010). By using the unpaired t-test, it is found that the cortical surface area of male is statistically larger than that of female on both hemispheres of all 3 ages, with the $P$ value consistently less than 0.01. By using the paired t-test, at 0-year of age, it is found that the surface area of the left hemisphere is statistically ($P<0.001$) larger than that of the right hemisphere, consistent to the report stating that the volume of the left hemisphere is larger than that of the right hemisphere at birth (Gilmore et al. 2007), although no difference is found in hemispheric surface area due to small sample in (Hill, Dierker et al. 2010). However, no significant difference on the cortical surface areas between the left and right hemispheres has been found at 1 and 2 years of age, consistent to the findings on adults (Van Essen et al. 2011), where nearly same cortical surface areas of left versus right hemispheres are observed.

Table 2 provides the mean and standard deviation of the longitudinal cortical surface area expansion of the 73 subjects from 0 to 2 years of age. The surface area expansion is defined as the ratio of the later year central cortical surface area versus that of the earlier year. In the first year, the cortical surface area expands dramatically, with the average expansion around 1.80 times. By using paired t-test, it is found that the cortical surface area expansion of the right hemisphere is statistically ($P<0.001$) larger than that of the left hemisphere from 0 to 1 year of age. This might explain the above findings: although the cortical surface area of the left hemisphere is statistically larger than that of the right hemisphere at 0 year of age, no significant difference between the cortical surface areas of left and right hemispheres exists at 1 year of age. In the second year, the cortical surface area expansion is still substantial, with the average expansion around 1.20 times. No significant difference on the cortical surface area expansion between the left and right hemispheres has been found from 1 to 2 years of age. Our results are consistent with the results in (Knickmeyer et al. 2008; Gilmore et al. 2011), where cortical gray matter volume growth is rapid in the first 2 years of life, especially in the first year. Although the cortical surface area of male is statistically larger than that of female in each hemisphere at each age, no significant difference in the cortical surface area expansion between male and female has been found from 0 to 1 year of age and from 1 to 2 years of age.

Longitudinal Surface-Based Atlases at 0, 1, and 2 Years of Age

Figure 3 shows the generated longitudinal surface-based atlases of cortical structures of the left hemisphere at 0, 1, and 2 years of age. Major cortical folding patterns are similar in terms of average convexity and mean curvature at 0, 1, and 2 years of age, suggesting that the shapes of major cortical folding of sulci and gyri have been well developed at term birth, which is consistent to the findings in (Hill, Dierker et al. 2010). However, no significant difference on the cortical surface areas between the left and right hemispheres has been found at 1 and 2 years of age, consistent to the findings on adults (Van Essen et al. 2011), where nearly same cortical surface areas of left versus right hemispheres are observed.
et al. 2010), and are well preserved during cortical development from 0 to 2 years of age. However, some minor differences on the appearance of cortical folding can still be observed, such as the increasing trend of the magnitude of average convexity and the decreasing trend of the magnitude of mean curvature from 0 to 2 years of age. The same findings have also been observed on the right hemisphere.

Figure 4 shows the average central cortical surfaces of the 73 subjects at 0, 1, and 2 years of age, generated by averaging the 3D positions of corresponding vertices of all cortical surfaces after affine alignment to the age-matched infant image intensity atlas (Shi et al. 2011). In the first year, the average cortical surface expands dramatically, while, in the second year, the average cortical surface expands much less than the
first year, although the expansion is still considerable. Meanwhile, the distance from the gyral crest to the sulcal bottom increases considerably from 0 to 2 years of age. However, the shapes of major cortical folding of sulci and gyri are quite similar at 0, 1, and 2 years of age, consistent with the above finding of cortical folding measurements of average convexity and mean curvature in Figure 3.

Region-Specific Surface Area Expansion in the First Year

Figure 5a shows the map of longitudinally cortical surface area expansion on both hemispheres from 0 to 1 year of age. Each region of the cortical surface increases in absolute surface area; however, the areal expansion is highly nonuniform across different cortical regions. In general, the areal expansions in the parietal, temporal, and occipital lobes are larger than those of the frontal lobes in both hemispheres. Specifically, high-expansion regions are concentrated in superior and medial temporal cortices, superior parietal cortices, medial orbitofrontal cortices, lateral anterior prefrontal cortices, occipital lobes, and postcentral gyri in both hemispheres. Low-expansion regions are concentrated in precentral gyri, superior frontal cortices, orbitofrontal cortices, cingulate cortices, precuneus regions, and insular regions in both hemispheres. By using t-tests with setting of significant level as $P \leq 0.01$ (corrected for multiple comparisons with 5000 iterations of permutation testing [Nichols and Holmes 2002]), the white contours in the Figure 5a enclose the cortical regions that expand significantly larger than the average expansion, whereas the black contours enclose the cortical regions that expand significantly smaller than the average expansion. In general, the regions passing significance are similar in the 2 hemispheres.

Region-Specific Surface Area Expansion in the Second Year

Figure 5b shows the map of longitudinally cortical surface area expansion from 1 to 2 years of age. Similar to the finding in the first year, each region of the cortical surface increases in absolute surface area, while the expansion is also highly nonuniform across different cortical regions. However, high-expansion and low-expansion regions from 1 to 2 years of age are quite different from those from 0 to 1 year of age. In general, the areal expansions in the parietal, temporal, and frontal lobes are larger than those of the occipital lobes in both hemispheres, consistent to the findings that the gray matter volumes of parietal lobes, frontal lobes, and temporal poles grow relatively fast (Gilmore et al. 2011) in the second year. Specifically, the high-expansion regions are concentrated in superior and middle frontal cortices, orbitofrontal cortices, inferior temporal cortices, inferior parietal cortices, superior parietal cortices, and isthmus of the cingulate cortex in both hemispheres. The low-expansion regions are concentrated in central sulci, paracentral lobules, superior temporal cortices, occipital lobes, and insular regions in both hemispheres. Similarly, the white (or black) contours enclose the cortical regions that expand significantly larger (or smaller) than the average areal expansion, respectively. In general, the regions passing significance are similar in both hemispheres.

Discussion

For the first time, we have quantitatively characterized the pattern of longitudinal cortical surface area expansion in the first 2 years of life by establishing the first longitudinal surface-based atlas of human cortical structures at 0, 1, and 2 years of age, which will facilitate the surface-based analysis of cortical folding and development in early postnatal stages of life. We find that major cortical folding of sulci and gyri
have been well developed at term birth, consistent with previous findings (Retzius 1890; Chi et al. 1977; Hill, Dierker et al. 2010), and are well preserved during cortical development from 0 to 2 years of age. Only the tertiary folding structures are still undergoing rapid development after birth. Therefore, the considerable variability of cortical folding across adult individuals is primarily established by term birth (Hill, Dierker et al. 2010). Cortical connectivity is thought as the major driving force of the cortical folding (Van Essen 1997; Nie, Guo et al. 2011), where consistent cortical folding is predicted to occur in regions with cortical connections dominated by few pathways between large areas (Van Essen 1997; Hill, Dierker et al. 2010). Since at term birth nearly every long-distance cortico-cortical and callosal connection has formed and only short cortico-cortical connections are still developing (Kostovic and Jovanov-Milosevic 2006; Takahashi et al. 2012), our result supports the hypothesis that cortical connectivity drives the variability of cortical folding (Van Essen 1997; Nie, Guo et al. 2011). Cortical folding abnormalities, which have been found in many neurodevelopmental disorders, such as autism (Nordahl et al. 2007), Williams syndrome (Van Essen et al. 2006), and bipolar disorder (Fornito

Figure 5. The maps of longitudinally cortical surface area expansion from 0 to 2 years of age. The color bar is shown on the right. The white and black contours enclose the cortical regions that expand significantly larger and smaller than the average area expansion, respectively.
et al. 2007), might have origins at term birth. Therefore, studying the cortical folding and development of neonates with high risks of neurodevelopmental disorders would help understand these disorders.

We find that the cortical surface area expansion is age related and region specific. In the first year, the surface area expands 1.80 times and the high-expansion regions are concentrated in superior temporal (auditory cortex), superior parietal (involved with multisensory integration [Molholm et al. 2006]), medial orbitofrontal, and lateral anterior prefrontal cortices, as well as postcentral gyrus (primary somatosensory cortex) and occipital lobe (visual cortex). This pattern might reflect the rapid development of visual, auditory, and sensory functions relative to the motor and association areas in the first year. In the second year, the surface area expands 1.20 times and the high-expansion regions are concentrated in the superior frontal (involved with motor planning [Schilling et al. 2012]), middle frontal (involved with self-evaluation [Beer et al. 2010]), inferior temporal (involved with high-order visual processing, including shapes and faces [Denys et al. 2004; Sabatinielli et al. 2011]), superior parietal (involved with visuospatial and attentional processing [Nachev and Husain 2006]), orbitofrontal (involved with decision making [Kringelbach 2005]), and inferior parietal cortices (involved with receiving auditory, visual, and somatosensory inputs [Roazzi et al. 2008]).

Cortical neurogenesis and migration are completed by the first week of postnatal life (Rakic 2009; Clowry et al. 2010; Hill, Inder et al. 2010). After that, cortical development is largely dependent on dendritic growth, growth of the terminal axon arborization, myelination, and synaptogenesis (Flechsig et al. 1896; Mrzljak et al. 1990; Koenderink et al. 1994; Rakic et al. 1994; Koenderink and Uylings 1995; Petanjek et al. 2011). Dendritic outgrowth during the first 2 years might count more for changes in cortical volume (Petenjak et al. 2008). In prefrontal layer IIIC pyramidal neurons, the total number of dendritic segments reached adult levels at the age of 1 postnatal month, and the total length of basal dendritic tree increased 3 times from birth to 2.5 months, remained “dormant” between 2.5 and 16 months and increased again until 2.5 years (Petenjak et al. 2008). In prefrontal layer V pyramidal neurons, no new dendritic segments were elaborated after birth, and the values of dendritic length became close to adult values at 12–15 months, with the most intensive period of postnatal dendritic growth occurring during the first 3 postnatal months (Petenjak et al. 2008). The basic unit of the mature neocortex is the minicolumn, a narrow chain of neurons extending vertically across the cellular layers II–VI, perpendicular to the pial surface (Mountcastle 1997). At the end of gestation, the mean width of a minicolumn was about one-third adult size (Buxhoeveden and Casanova 2002), and the neurons are rarely generated after birth in the cerebral cortex (Rakic 1988). Therefore, the expansion of cortical surface area is likely caused by expansions of minicolumns.

Human brain is regionally highly differentiated but hierarchically and geometrically organized (Chen et al. 2012; Wedeen et al. 2012; Zilles and Amunts 2012). The cortical subdivision are nested in a common superstructure and interrelated in a tree-like organization, showing different degrees of functional, molecular, genetic, or structural similarities. The region-specific cortical surface area expansion might relate to cellular, functional, and genetic nonuniformities during development. In general, high-expansion regions in the first year tend to peak synaptic density (Huttenlocher and Dabholkar 1997), gray matter density (Gogtay et al. 2004), cortical thickness (Shaw et al. 2008), and functional maturation (Chugani and Phelps 1986; Chugani 1998) earlier than the high-expansion regions in the second year. The longitudinal regional variation of cortical surface area expansion supports the regional differences in synapse development in human brain development, with the synaptic density peaked at 3 months in the auditory cortex and at 3.5 years in the middle frontal gyrus (Huttenlocher and Dabholkar 1997). Similarly, the somatic sensory cortex (postcentral gyrus) and occipital pole peak cortical thickness earlier than high-order cortical areas, such as prefrontal and cingulate cortices (Shaw et al. 2008). And the sensorimotor cortex and occipital pole peak gray matter density earlier than the regions involved in executive function, attention, and motor coordination (prefrontal cortex) (Gogtay et al. 2004). Motor, visual, auditory, and somatosensory cortices are the early myelinating regions (Flechsig et al. 1896). In the first 3 postnatal months, glucose metabolic activity in PET increases more in the parietal, temporal, and primary visual cortices than the frontal cortex (Chugani and Phelps 1986; Chugani 1998). In the first 2 postnatal months, PET imaging or scalp recording can detect the activity in some high-expansion regions in the first year, including auditory cortex in response to sound and the medial temporal cortex in response to visually presented faces (de Haan and Nelson 1997; Tzourio-Mazoyer et al. 2002; Wakai et al. 2007; Hill, Inder et al. 2010). In contrast, transcranial magnetic stimulation of the motor cortex does not elicit detectable muscle response until 2 years of age (Nezu et al. 1997; Martin 2005; Hill, Inder et al. 2010). This longitudinal cortical surface area expansion pattern is also consistent with the cortical functional network development. For example, in infants, the resting-state networks are present in sensorimotor, visual, and auditory cortices (Fransson et al. 2007; Lin et al. 2008), with cortical hubs found in motor, sensory, auditory, and primary visual cortices (Fransson et al. 2011). However, the default network involving inferior parietal, prefrontal, and temporal cortices does not fully develop until 2 years of life (Gao et al. 2009).
unrelated to the regions with larger cortical surface area expansion in the first year, suggesting that the cortical volume growth is jointly contributed by both cortical surface area expansion and cortical thickness growth at this stage. These findings may have clinical implications for understanding abnormal cortical development in neurodevelopmental disorders. For example, autism is associated with overgrowth of cortical volume in the first 2 years of life (Schumann et al. 2010; Hazlett et al. 2011), which are primarily caused by the increase of the cortical surface area rather than the cortical thickness (Hazlett et al. 2011). Since the cortical surface area expansion in the first year is much larger than that of the second year, the cortical volume overgrowth in autism might occur more likely in the first year (Gilmore et al. 2011). By correlating the longitudinal cortical surface expansion patterns with the adult cortical thickness map (Fischl and Dale 2000), the high-expansion regions in the first year (i.e., post-central gyrus, superior parietal cortex, and occipital cortex) tend to be relatively thin in adults. This correlation is imperfect with the noteworthy exceptions including high-expansion regions of superior temporal, lateral anterior prefrontal, and medial orbitofrontal cortices, which are relatively thick in adults. All these implications and questions need to be further confirmed and answered with a longitudinal study of cortical thickness development in the first 2 years of life.

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